US ERA ARCHIVE DOCUMENT

CASWELL FILE

007871



UNITED STATES ENVIRONMENTAL PROTECTION AGENCY WASHINGTON, D.C. 20460

APR 23 1990

MEMOR ANDUM

SUBJECT:

CYPROCONAZOLE - DOG CHRONIC FEEDING AND RAT CHRONIC FEEDING/ONCOGENICITY STUDIES

SUSAN LEWIS/GRABLE TO:

PRODUCT MANAGER (21)

REGISTRATION DIVISION (H7505C)

FR OM:

LINDA L. TAYLOR, PH.D.

TOXICOLOGY BRANCH II, SECTION IT

HEALTH EFFECTS DIVISION (H7509C)

THRU

K. CLARK SWENTZEL / LUCION SECTION II HEAD, TOXICOLOGY BRANCH II

HEALTH EFFECTS DIVISION (H7509C)

AND

muan Emect 4/19/90 MARCIA VAN GEMERT, PH.D. MURLY XXIII (H7509C)

REGISTRANT:

SANDOZ CROP PROTECTION CORPORATION

CHEMICAL:

CYPROCONAZOLE SAN 619 F

SYNONYMS: PROJECT:

9-2056

CASWELL NO .: MRID No 👀

412129-01 & 411647-01

ECORD No .:

NONE PROVIDED

DENTIFYING NO .:

55947-RGG

ACTION REQUESTED:

NONE. Two CHRONIC STUDIES SUBMITTED.

IN A LETTER DATED JULY 10, 1989, THE REGISTRANT SUBMITTED THE TWO STUDIES REFERENCED ABOVE, IN ORDER TO PROVIDE THE AGENCY WITH A COMPLETE PICTURE OF THE CHRONIC TOXICOLOGY PACKAGE FOR SAN 619 F. A MOUSE ONCOGENICITY STUDY (MR ID # 411472-01) WAS SUBMITTED UNDER SEPARATE COVER "FYI". THE DER FOR EACH (DOG AND RAT) STUDY IS ATTACHED.

<u>Dog study</u> - The administration of SAN 619 F to beagle dogs for 52 weeks at dose levels of 30, 100, and 350 ppm [1.0, 3.2, and 12.1 (males); 12.6 (females) MG/KG/DAY, RESPECTIVELY RESULTED IN DIFFERENCES IN SEVERAL CLINICAL LABORATORY PARAMETERS BETWEEN THE CONTROL AND TREATED ANIMALS. ABSOLUTE AND RELATIVE LIVER WEIGHTS WERE INCREASED IN THE HIGH-DOSE ANIMALS OF BOTH SEXES COMPARED TO CONTROLS, BUT STATISTICAL SIGNIFICANCE WAS ATTAINED ONLY IN THE MALES. RELATIVE KIDNEY WEIGHT WAS INCREASED (SIGNIFICANTLY) IN BOTH THE LOW- AND HIGH-DOSE FEMALES. THE NOEL CAN BE SET AT 30 PPM (1.0 MG/KG/DAY) AND THE LEL AT 100 PPM (3.2 MG/KG/DAY), BASED ON LIVER EFFECTS.

CLASSIFICATION: CORE MINIMUM UNDER GUIDELINE 83-1, CHRONIC TOXICITY.

Printed on Recycled Paper

RAT STUDY - THE ADMINISTRATION OF SAN 619 F TO MALE RATS FOR 118 WEEKS AND TO FEMALE RATS FOR 121 WEEKS AT DOSE LEVELS OF 0, 20, 50, OR 350 PPM (MALES - 1.0, 2.2, AND 15.6 MG/KG; FEMALES - 1.2, 2.7, 21.8 MG/KG) RESULTED IN DECREASED BODY WEIGHTS IN HIGH-DOSE FEMALES AND INCREASED INCIDENCE OF FATTY INFILTRATION OF THE LIVER IN THE HIGH-DOSE MALES. THE NOEL FOR SYSTEMIC TOXICITY CAN BE SET AT 50 PPM, THE LEL AT 350 PPM, BASED ON DECREASED BODY WEIGHT IN FEMALES AND FATTY INFILTRATION IN THE LIVER OF MALES.

Under the conditions of the study, SAN 619 F was not carcinogenic. Based on the Lack of: 1) any biologically significant body weight decrement; 2) any significant histopathological correlate accompanying the increased relative liver weight; 3) any increase in the liver enzyme activities in the females; 4) any consistent change in the liver enzyme activities in the high-dose males, suggests that the dose levels chosen were not adequate to determine the carcinogenic potential of the test material.

CLASSIFICATION: CORE SUPPLEMENTARY UNDER GUIDELINE 83-2, CARCINOGENICITY STUDY; CORE MINIMUM UNDER GUIDELINE 83-1, CHRONIC TOXICITY STUDY.

REVIEWED BY: LINDA L. TAYLOR, PH.D. MAR TOX. BRANCH II, SECTION II (H75096)

SECONDARY REVIEWER: K. CLARK SWENTZEL A Clark HEAD SECTION II, Tox. BRANCH II (H7509C)

DATA EVALUATION REPORT

STUDY TYPE: ONE-YEAR CHRONIC - DOG

272E TUX. CHEM. NO.:

MR ID NO -: 412129-01

ALPHA-(4-CHLOROPHENYL)-ALPHA-(1-CYCLOPROPYLETHYL)-1H-TEST MATERIAL:

1,2,4-TRIAZOLE-1-ETHANOL; SAN 619F

S YNONYMS: CYPROCONAZOLE

PROJECT No. 394-D STUDY NUMBER:

SPONSOR: SANDOZ CROP PROTECTION CORPORATION

TESTING FACILITY: SANDOZ LTD. AGRO DEVELOPMENT - TOXICOLOGY DEPARTMENT

BASLE/SWITZERLAND

CHRONIC URAL TOXICITY BY DIETARY ADMINISTRATION TO BEAGLE TITLE OF REPORT:

DOGS FOR ONE YEAR

AUTHOR(S): S.F.P. WARREN, F. HAMBURGER, S. CARPY, AND F. MILLER

REPORT ISSUED: April 18, 1988

QUALITY ASSURANCE: A QUALITY ASSURANCE STATEMENT WAS PROVIDED.

CLASSIFICATION: CORE-MINIMUM.

CUNCLUSION: THE ADMINISTRATION (DIET) OF SAN 619 F TO BEAGLE DOGS FOR 52 WEEKS AT DOSE LEVELS OF 30, 100, AND 350 PPM [1.0, 3.2, AND 12.1 (MALES); 12.6 (FEMALES) MG/KG/DAY, RESPECTIVELY RESULTED IN DIFFERENCES IN SEVERAL CLINICAL LABORATORY PARAMETERS BETWEEN THE CONTROL AND TREATED ANIMALS, WHICH ARE CONSISTENT WITH EFFECTS ON THE LIVER (ELEVATED ALKALINE PHOSPHATASE AND ALAT LEVELS; DECREASED TOTAL PROTEIN, ALBUMIN, AND CHOLESTEROL LEVELS). ABSOLUTE AND RELATIVE LIVER WEIGHTS WERE INCREASED IN THE HIGH-DOSE ANIMALS OF BOTH SEXES COMPARED TO CONTROLS, BUT STATISTICAL SIGNIFICANCE WAS ATTAINED ONLY IN THE MALES. RELATIVE KIDNEY WEIGHT WAS INCREASED (SIGNIFICANTLY) IN BOTH THE LOW- AND HIGH-DOSE FEMALES.

STATISTICALLY SIGNIFICANT INCREASES WERE OBSERVED IN CYTOCHROME P450 IN BOTH SEXES OF THE HIGH-DOSE AND IN THE MID-DOSE FEMALES. LAMINAR EOSINOPHILIC INTRAHEPATOCYTIC BODIES WERE OBSERVED IN ALL HIGH-DOSE MALES, ONE MID-DOSE MALE, AND TWO HIGH-DOSE FEMALES AND WERE CONSIDERED TO REPRESENT ADAPTIVE HYPERTROPHY OF THE ENDOPLASMIC RETICULUM.

THE NUEL CAN BE SET AT 30 PPM (1.0 MG/KG/DAY) AND THE LEL AT 100 PPM (3.2 MG/KG/DAY), BASED ON LIVER EFFECTS.

A. MATER IALS:

- 1. Test compound: Sandoz 619F, technical; Description: A Light Brown Powder; Batch # 8507; Purity: 95+ 1%; Stability: Documented.
- 2. Test animals: Species: Dog; Strain: purebred Beagles (Canis Familiaris); Age: approximately 5 months; Weight: males: 7.6-8.0 kg, Females: 7.0-7.5 kg; Source: Marshall Farms Breeding Laboratories, North Rose, New York.

B. STUDY DESIGN

1. ANIMAL ASSIGNMENT

ANIMALS WERE ASSIGNED RANDOMLY TO THE FOLLOWING TEST GROUPS:

GROUP	DOSE IN DIET (PPM)	Exposure Period (MALES	(52 MONTHS) FEMALES	
1 CONTROL	0	4	4	
2 LOW	30	4	4	
3 MID	100	4	4	
4 HIGH	350	4	4	

2. DIET PREPARATION

THE TEST ARTICLE WAS ADMINISTERED IN THE DIET (POWDERED DIET # 24-335-1; KLINGENTALMUHLE, CH-BASLE). A DIET PREMIX WAS PREPARED MONTHLY [40 GRAMS OF SAN 619F MIXED WITH 3960 GRAMS OF POWDERED FODDER CONCENTRATION OF 10 MG SAN 619F PER GRAM (1%)] WHICH WAS THEN MIXED WEEKLY BY ADDING ADDITIONAL POWDERED FODDER TO MAKE THE APPROPRIATE DOSE LEVELS. THE CONTROLS RECEIVED UNTREATED DIET. EACH DOG WAS OFFERED APPROXIMATELY 400 GRAMS OF FOOD DAILY AND WATER AD LIBITUM.

RESULTS

Data provided indicate the diets to be within +/-10% of the target concentrations throughout the study-

3. STATISTICS - THE PROCEDURES UTILIZED IN ANALYZING THE NUMERICAL DATA ARE EXPLAINED ON PAGE 17 OF THE FINAL REPORT (COPY ATTACHED).

C. METHODS AND RESULTS:

1. OBSERVATIONS

ALL DOGS WERE OBSERVED TWICE DAILY FOR SIGNS OF ILL HEALTH, AND THE NOSE, EYES, ANUS, AND BUCCAL CAVITY WERE CHECKED DAILY.

RESULTS:

TOXICITY/MORTALITY (SURVIVAL)

THERE WERE NO DEATHS OR UNSCHEDULED SACRIFICES DURING THE STUDY.

CLINICAL OBSERVATIONS

SUBDUED BEHAVIOR (DURING WEEKS 4 TO 8) AND LOWER BODY-WEIGHT GAIN (DURING THE FIRST 4 WEEKS) WERE OBSERVED IN ONE HIGH-DOSE FEMALE. FOOD CONSUMPTION ALSO TENDED TO BE LOW IN THIS ANIMAL UP TO WEEK 10. THEREAFTER, THIS ANIMALS DISPLAYED COMPARABLE BEHAVIOR TO THE OTHER DOGS. ONE MID-DOSE FEMALE DISPLAYED SUBDUED BEHAVIOR WITH BODY WEIGHT LOSS AND LOW FOOD CONSUMPTION DURING WEEK 2.

ONE HIGH-DOSE FEMALE FAILED TO COME INTO HEAT DURING THE STUDY, AND TERMINAL HISTOLOGICAL EXAMINATION REVEALED IMMATURE OVARIES.

2. BODY WEIGHT AND FOOD CONSUMPTION

ANIMALS WERE WEIGHED WEEKLY AND FOOD CONSUMPTION WAS DETERMINED WEEKLY. GROUP MEAN WEEKLY INTAKE OF TEST MATERIAL WAS CALCULATED FROM INDIVIDUAL BODY WEIGHT AND FOOD CONSUMPTION.

RESULTS

THE AUTHOR REPORTED AN INITIAL EFFECT ON BODY-WEIGHT GAIN FOR THE HIGH-DOSE MALES; A DECREASE WAS NOTED INITIALLY, WHICH WAS GREATEST AT WEEK 9 AND THEN BECAME COMPARABLE TO CONTROL THEREAFTER.

			Вс	DY WEI	GHT (%	CONTR	OL)			·
WEEK	0	1	3	6	9	14	15	16	26	50
MALES LOW MID HIGH	96 101 100	100 104 100	100 105 99	97 103 95	97 103 94	97 103 93	98 104 93	98 105 93	102 104 97	117 114 105
WEEK	0	1	3	6	9	14	27	44	47	50
FEMALES LOW MID HIGH	103 101 96	100 101 93	101 100 94	98 100 90	100 107 95	100 111 98	104 107 99	98 107 93	95 104 92	101 111 93

NOTE: DUE TO A COMPUTER PROBLEM AT THE TESTING FACILITY, PRE-TREATMENT VALUES FOR BOTH BODY WEIGHT AND FOOD CONSUMPTION WERE NOT AVAILABLE.

BODY-WEIGHT GAIN (KG)

- MALES	WEEK 0 - 9	WEEK 0 - 13	WEEK 13 - 26	WEEK 26 - 52	week 0 - 52
C L M H	1.9 1.9 2.1 1.3	2.3 2.5 2.7 1.8	1·1 1·4 1·1 1·3	-0·1 1·5 0·7 0·5	3.3 5.4 4.5 3.6
FEMALES C L M H	1.0 (1.0) 0.8 (0.6) 1.5 (0.9) 0.9 (0.5)	* 1.1 0.9 1.9 1.1	0.6 0.6 0.4 0.7	0.5 0.5 0.9 0.0	2·2 2·0 3·2 1·8

 $^{^{\}star}$ value for body-weight gain between weeks 0 and 6

In general, mean food consumption of all of the male groups was significantly greater than control at various time points during the study. For females, food consumption varied considerably, with the high-dose group showing decreased intake, compared to the controls and other treated groups for the first 3 weeks of the study and during weeks 31, 36, 42, 47, and 51 (not statistically significant). The low-dose females showed decreased food consumption from week 23 to week 48 (statistically significant at week 26 only), and the mid dose showed a decrease at weeks 31, 36, 37, and during weeks 42-47 (none statistically significant but $\geq 10\%$ below control value).

FOOD EFFICIENCY DATA PROVIDED IN THE REPORT INDICATE THAT THERE WAS A SLIGHTLY REDUCED EFFICIENCY AT THE HIGH DOSE FOR MALES OVER THE FIRST 26 WEEKS. FOR THE FIRST 9 WEEKS, BOTH SEXES AT THE HIGH DOSE DISPLAYED A REDUCED FOOD EFFICIENCY (NO STATISTICS WERE PERFORMED ON THESE DATA).

3. COMPOUND INTAKE

AVERAGE DAILY DOSES OF THE TEST MATERIAL WERE CALCULATED FROM NOMINAL DIETARY CONCENTRATIONS, INDIVIDUAL BODY WEIGHT, AND FOOD CONSUMPTION.

	CONSUMED	(MG/KG/DAY)
	MALES	FEMALES.
Low	0.99	0.99
MID	3.15	3.23
HI GH	12.05	12.58

4. UPHTHALMOLOGICAL EXAMINATIONS

OCULAR EXAMINATIONS WERE PERFORMED ON ALL OF THE DOGS PRIOR TO TREATMENT AND DURING WEEK 50, USING A FUNDUS CAMERA FOLLOWING ADMINISTRATION OF A MYDRIATIC.

6

RESULTS

THERE WERE NO TREATMENT-RELATED OCULAR CHANGES IN EITHER SEX.

5. BLOOD ANALYSIS

CLINICAL LABORATORY STUDIES WERE CONDUCTED ON ALL DOGS (FASTED 16-18 HOURS) PRIOR TO STUDY INITIATION (WEEKS -1 & -2), AT 3, 6, AND 9 MONTHS, AND AT STUDY TERMINATION. THE CHECKED (X) PARAMETERS WERE EXAMINED.

X ERYTHROCYTE COUNT (RBC)* X PLATELET COUNT* THROMBOCYTE COUNT X PROTHROMBIN TIME X PROTHROMBIN TIME X MEAN CORPUSCULAR VOLUME (MCV) X RETICULOCYTE COUNT METHEMOGLOBIN HEINZ BODIES	PLATELET COUNT* THROMBOCYTE COUNT PROTHROMBIN TIME	X X X X	RETICULOCYTE COUNT METHEMOGLOBIN HEINZ BODIES	;)
ERYTHROCYTE INDICES HOWELL-JOLLY BODIES			HOWELL-JOLLY BODIES	

^{*} REQUIRED FOR CHRONIC STUDIES

RESULTS

Increased platelet counts were observed at the high dose in both sexes throughout the study. Mid-dose males also displayed marginally increased counts at weeks 26, 38, and 52. Prothrombin time was measured at week 48 to assess whether the disturbed platelet count was associated with disturbed clotting function. The increase in prothrombin time observed was considered (by the author) to be the result of the occasional low values in the low-dose and control groups, and not to treatment. However, it is to be noted that the low- and mid-dose females showed lower values than the control females.

HEMATOLOGICAL VALUES (% OF CONTROL)

PAR AMETER							
MALES	OR OUP	WEEK -1	WEEK 13	WEEK 26	WEEK 38	WEEK 48	WEEK 52
PLATELETS	CONTROL	403	305	287	269	291	274
TH/CMM	LOW	334(83)	269(88)	256(89)	248(92)	266(91)	258(94)
,	MID	386(96)	335(110)	356(124)	307(114)	332(114)	318(116)
	HIGH	358(89)	461(151)	423(147)	384(143)	409(140)	383(140)
PROTHROMB •		220(03)	-101(1)1/	122(11)	JU 1(1 1)	9-60	<u> </u>
SEC •	LOW					10.50(109)	
JEC.						10.80(113)	
	MID					10.85(113)	
FEMALES.	HIGH	UECV O	UCCV 17	VICEN OC	UCEL ZO		UCEIZ CO
FEMALES	GR OUP	WEEK O	WEEK 13	WEEK 26	WEEK 38	WEEK 48	WEEK 52
PLATELETS	CONTROL	374	319	324	323	310	305
TH/CMM	LOW	352(94)	291(91)	317(98)	308(95)	322(104)	324(106)
	MID	362(97)	344(108)	351(108)	328(102)	371(120)	332(109)
	HIGH	344(92)	456(143)*	433(134)	415(129)	442(143)*	<u>438(144)**</u>
PROTHROMB .						10.08	•
SEC.	LOW					9.88(98)	
	MID					9.83(98)	
	HI GH					11.05(110)	
							

B. CLINICAL CHEMISTRY

THE CHECKED (X) PARAMETERS WERE MEASURED.

ELECTROLYTES: X CALCIUM* X CHLORIDE* MAGNESIUM* X PHOSPHORUS* X POTASSIUM* X SODIUM* ENZYMES X ALKALINE PHOSPHATASE CHOLINESTERASE# X CREATININE PHOSPHOKINASE* X LACTIC ACID DEHYDROGENASE X SERUM ALANINE AMINOTRANSFERASE X SERUM ASPARTATE AMINOTRANSFERASE GAMMA GLUTAMYL TRANSPEPTIDASE	ASE (ALSO SGOT)*
X GLUTAMATE DEHYDROGENASE	A LEOCINE ARTLAMIDASE

* REQUIRED FOR CHRONIC STUDIES # SHOULD BE REQUIRED FOR OP

RESULTS

THERE WAS A STATISTICALLY SIGNIFICANT DECREASE IN TOTAL PROTEIN, ALBUMIN, AND CHOLESTEROL VALUES AT VARIOUS TIME POINTS THROUGHOUT THE STUDY IN BOTH SEXES AT THE HIGH-DOSE. TOTAL BILIRUBIN VALUES ALSO TENDED TO BE DEPRESSED, ESPECIALLY IN THE HIGH-DOSE FEMALES, BUT STATISTICAL SIGNIFICANCE WAS NOT ATTAINED. ALKALINE PHOSPHATASE VALUES WERE INCREASED IN BOTH SEXES OF THE HIGH DOSE THROUGHOUT THE STUDY (STATISTICALLY SIGNIFICANT AT ALL TIME POINTS IN THE FEMALE, BUT ONLY AT WEEK 51 FOR MALES). THE MID-DOSE FEMALES DISPLAYED A SIMILAR INCREASE OF LESSER MAGNITUDE, BUT STATISTICAL SIGNIFICANCE WAS NOT ATTAINED. THE MID-DOSE MALES DISPLAYED AN INCREASE ONLY AT THE 51-WEEK TIME POINT, WHICH WAS NOT STATISTICALLY SIGNIFICANT. CALCIUM LEVELS WERE DECREASED (P<0.01) AT THE 13-WEEK TIME POINT IN BOTH SEXES OF THE HIGH DOSE AND AT THE 25-WEEK TIME POINT IN THE HIGH-DOSE MALES. ALAT VALUES FOR THE HIGH-DOSE MALES WERE ELEVATED AT ALL TIME POINTS DURING THE STUDY COMPARED WITH CONTROL VALUES, BUT STATISTICAL SIGNIFICANCE WAS ATTAINED ONLY AT WEEK 51. SIMILAR EFFECTS WERE NOT OBSERVED IN THE FEMALES. GLOB % was elevated significantly in the high-dose (both sexes) at weeks 13 AND 25, AND IN THE HIGH-DOSE MALES AT WEEK 38. TRIGLYCERIDE VALUES WERE SIGNIFICANTLY DEPRESSED IN THE HIGH-DOSE FEMALE AT WEEKS 13 AND 38.



CLINICAL CHEMISTRY VALUES (% OF CONTROL)

PAR AMETER	GR OUP	WEEK -1	WEEK 13	WEEK 25	WEEK 38 70	WEEK 51 67
MALES T. Protein	CONTROL	59 58(98)	65 63(97)	69 67(98)	68(97)	66(99)
I. PROTEIN G/L	LOW MID	58(98)	62(96)	69(101)	67(95)	66(98)
	HI GH	58(99)	56(86)**	61(89)**	59(85)**	62(92)
ALBUMIN	CONTROL	25.7	28.6	31.3	30.1	32.9
G/L	LOW	25.7(100)	27.8(97)	30.1(96)	28.4(94)	31.0(94)
	MID	25.3(98)	26-9(94)	29.0(93)	27·0(90) 20·2(67)**	30•3(92) 25•6(78)**
CHOL FOTEROL	HIGH	23.6(92) 2.89	21·2(76)** 3·82	23.6(76)** 3.29	3.48	3.59
CHOLESTEROL MMUL/L	CONTROL LOW	2.66(92)	2.96(78)	2.93(89)	3·04(87)	3.38(94)
MINUL/ L	MID	2.39(83)	2.76(72)	2.80(85)	2.56(74)	3.05(85)
	HIGH	2.37(82)	1.38(36)**	1.80(55)	1.72(49)*	2-07(58)
BILIRUBIN	CONTROL	1.453	1.557	1.570	1.608	1.648
UMOL/L	LOW	1.495(103)	1.403(90)	1.648(105)	2.140(133)	1.708(104)
	MID	1.413(97)	1.420(91)	0.918(58)	1.713(107)	1.570(95)
	HI GH	1.565(108)	1.185(76)	1.500(96)	1.300(81)	1.340(81) 96.1
ALKALINE	CONTROL	403.3	298.0	137·8 105·4(76)	126•6 107•4(85)	80.5(84)
PHOSPHATASE	LOW	273.0(65) 299.5(69)	169·0(57) 206·3(69)	140.8(102)	128.8(102)	117.4(122)
U/L	MID HIGH	268.0(61)	380.3(128)	287-8(209)	362.3(286)	375.5(391)**
ALAT	CONTROL	20.28	21.77	27.98	32.53	28.85
U/L	LOW	24.38(120)	25.15(116)	28.10(100)	29.48(91)	27•00(94)
0, 2	MID	21.03(104)	25.08(115)	27.43(98)	27•83(86)	27.13(94)
	HIGH	19.10(94)	38.58(177)	40.05(143)	46.85(144)	48.57(16)
CALCIUM	CONTROL	2.740	2.698	2.675	2.545	2.593
MMOL/L	LOW	2.725(99)	2.608(97)	2.543(95)	2.425(95)	2.505(97) 2.483(96)
	MID	2.628(96)	2•578(96) 2•408(89)**	2·528(95) 2·400(90)**	2•403(94) 2•353(92)	2.473(95)
GLOB• %	HI GH	2.673(98) 56.55	56.11	54.68	57.14	51.17
GLOB• %	CONTROL LOW	55.78(99)	55.82(99)	55.22(101)	58.22(102)	53.38(104)
/0	MID	56.26(99)	56.94(101)	58.23(106)	59.31(104)	54.06(106)
	HI GH	59.64(105)	62.14(111)*	61.50(112)*	65.99(115)*	58.56(114)
PAR AMETER	GR OUP	WEEK -1	WEEK 13	WEEK 25	WEEK 38	WEEK 51
<u>FEMALES</u>	CONTROL	58.4	62.3	63.9	72 65•5(91)	65•2 64•7(99)
T. PROTEIN	LOW	59.8(102)	62.3(100)	68.1(107)	02.2(80)	62.9(97)
G/L	MID	55.6(95)	59•4(95) 54•4(87)*	62•2(97) 59•1(92)	63.6(88) 58.3(81)	59.3(91)
A. BUMIN	HIGH	59.9(103) 27.7	29.4	30.2	30.1	59.3(91) 32.0
ALBUMIN G/L	CONTROL LOW	27 . 3(99)	30.4(103)	29.6(98)	29.8(99)	30.7(96)
U/ L	MID	24.6(89)	28•3(96)	29.6(98) 28.5(94)	27.9(93)	30.7(96) 29.7(93)
	HIGH	26.9(97)	21.5(73)**	24.3(80)**	23.6(78)*	26.4(83)**
CHOLESTEROL	CONTROL	3-03	4.23	4.11	4• <i>7</i> 5	5.35
MMOL/L	LOW	2.87(94)	3.49(82)	5.19(126)	3.53(74)	4.56(85)
	MID	2-44(80)	3.98(94)	3. 79(92)	4.79(101)	4.59(86)
•	HI GH	2.51(83)	2.05(48)**	2.76(67)	2.45(52)**	<u>2.80(52</u>)*

PAR AMETER	GR OUP	WEEK -1	WEEK 13	WEEK 25	WEEK 38	WEEK 51
BILIRUBIN	CONTROL	1.815	2.013	1.925	2.550	2.090
UMOL/L	LOW	1.743(96)	1.950(97)	1.620(84)	1.955(77)	1.860(89)
DI IOLI L	MID	1.673(92)	1.703(85)	1.320(69)	2.265(89)	1.803(86)
	HI GH	1.933(107)	1.368(68)	1.528(79)	1.455(57)	<u>1•393(67</u>)
ALKALINE	CONTROL	226.0	146.6	122.0	137.5	108.9
PHOSPHATASE	LOW	197.8(88)	139.8(95)	105-6(87)	118.7(86)	98.6(91)
U/L	MID	229.0(101)	216.8(148)	164.5(135)	158.5(115)	152-8(140)
0/ L	HI GH	248.3(110)	407.8(278)*	366.8(301)**	407 • 8(297)*	<u>342.8(31</u> 6)**
ALAT	CONTROL	22.48	17.75	19.58	21.65	20.88
Ú/L	LOW	21.08(94)	20.53(116)	16.30(83)	22.75(105)	17.80(85)
U/ L	MID	23.65(105)	20.90(118)	21.55(110)	22.65(105)	22.20(106)
	HI GH	22.23(99)	21.45(121)	22.13(113)	22.43(109)	<u>22.85(10</u> 9)
CALCIUM	CONTROL	2.628	2.665	2 • 465	2.323	2.540
MMOL/L	LOW	2.703(103)	2.635(99)	2.530(103)	2.348(94)	2.548(100)
1910474	MID	2.603(99)	2.630(99)	2.455(100)	2.308(94)	2.578(101)
•	HIGH	2.620(100)	2.455(92)**	2.318(94)	2.193(89)	2•365(9 <u>3</u>)
GLOB. %	CONTROL	52.65	52.78	52.74	57-25	50-72
W N	LOW	54.44(103)	51.20(97)	56.52(107)	54.41(95)	52.56(104)
/0	MID	55.79(106)	52.39(99)	54.21(103)	56.12(98)	52.63(104)
		55.18(105)	60.54(115)**	58.91(112)*	59.35(104)	55.43(109)
	HIGH	77-10(1077	しし・フィ(エエン)	<u> </u>		

^{*} P<0.05

6. URINALYSIS

URINE WAS COLLECTED (AFTER A 16-18 HOUR FAST) FROM ALL ANIMALS PRE-DOSE, AND AT 3, 6, AND 9 MONTHS, AND PRIOR TO STUDY TERMINATION. THE CHECKED (X) PARAMETERS WERE EXAMINED.

		APPEARANCE*	X	GLUCOSE"
		VOLUME*	X	KETONES*
1	χl	SPECIFIC GRAVITY*	X	BILIRUBIN*
	χl	РН	X	BLOOD*
	χl	SEDIMENT (MICROSCOPIC)*		NITRATE
	χl	PROTEIN*	X	UROBILINOGEN
		DSMOLALITY		

^{*} REQUIRED FOR CHRONIC STUDIES

RESULTS

THERE WERE SPORADIC STATISTICALLY SIGNIFICANT ALTERATIONS IN A FEW URINARY PARAMETERS COMPARED TO CONTROL VALUES, BUT NONE THAT COULD BE ATTRIBUTED TO TREATMENT. THE PH OF THE HIGH-DOSE MALES WAS MORE ALKALINE THAN THE CONTROL VALUE AT WEEKS 12, 25, AND 38, BUT STATISTICAL SIGNIFICANCE WAS NOT ATTAINED.

GROSS PATHOLOGY

ALL ANIMALS WERE SACRIFICED AT STUDY TERMINATION AND WERE SUBJECTED TO GROSS PATHOLOGICAL EXAMINATION. ALL DOGS WERE EXAMINED EXTERNALLY BOTH VISUALLY AND

BY PALPATION, WITH PARTICULAR ATTENTION BEING PAID TO THE EYES, NOSE, BUCCAL CAVITY, EXTERNAL GENITALIA, AND ANUS. A MACROSCOPIC EXAMINATION WAS PERFORMED AFTER OPENING ALL CAVITIES AND OBSERVING THE TISSUES IN SITU. THE FOLLOWING ORGANS WERE WEIGHED:

ADRENAL	LIVER	SPLEEN
BRAIN	OVARI ES	TESTES
KIDNEY	PITUITARY	THYROID WITH PARATHYROID

RESULTS

Two high-dose males displayed enlarged livers at necropsy, with pronounced lobular patterning. No other treatment-related findings were reported.

ABSOLUTE AND RELATIVE LIVER WEIGHT WERE SIGNIFICANTLY INCREASED IN THE HIGH-DOSE MALES COMPARED TO CONTROLS. HIGH-DOSE FEMALES ALSO DISPLAYED AN INCREASE IN LIVER WEIGHT (BOTH ABSOLUTE AND RELATIVE), BUT A P<0.05 WAS NOT ATTAINED.

ABSOLUTE KIDNEY WEIGHT WAS SLIGHTLY INCREASED IN THE LOW- AND HIGH-DOSE FEMALES, BUT STATISTICAL SIGNIFICANCE WAS NOT ATTAINED. RELATIVE KIDNEY WEIGHT WAS SIGNIFICANTLY INCREASED IN BOTH THE LOW- AND HIGH-DOSE FEMALE GROUPS.

ABSOLUTE THYROID WEIGHT OF THE FEMALES SHOWED A SLIGHT INCREASE WITH DOSE, BUT THIS WAS NOT DOSE-RELATED. RELATIVE THYROID WEIGHT INCREASED WITH INCREASING DOSE, BUT THIS DID NOT ATTAIN STATISTICAL SIGNIFICANCE.

THE PITUITARY WAS ALSO FOUND TO BE HEAVIER IN THE HIGH-DOSE ANIMALS COMPARED TO THE RESPECTIVE CONTROLS, BUT STATISTICAL SIGNIFICANCE WAS NOT ATTAINED.

ORGAN WEIGHTS

MM 50	LIV ABSOLUTE GRAMS	R RELATIVE	· · -	NEY RELATIVE	THYR ABSOLUTE GRAMS	OID RELATIVE %	PITUI ABSOLUTE GRAMS	TARY RELATIVE
MALES CONTROL LOW MID HIGH	251 -279 296 400**	2•34 2•26 2•47 3•64*	45.0 54.3 50.5 53.8	0.43 0.44 0.42 0.49	0•79 0•75 0•73 0•86	0.0073 0.0061 0.0061 0.0077	61.8 84.8 67.0 75.3	0.57 0.69 0.56 0.68
FEMALES CONTROL LOW MID HIGH	258 297 282 290	2•76 3•24 2•78 3•39	33.8 37.4 34.7 36.7	0.36 0.41* 0.36 0.43*	0.66 0.67 0.82 0.77	0.0070 0.0073 0.0080 0.0091	89.5 82.5 70.8 111.0	0.96 0.89 0.69 1.31

^{*} P<0.05 ** P<0.01

LIVER CHEMISTRY

AT NECROPSY, LIVER SAMPLES WERE TAKEN FROM ALL DOGS FOR ANALYSIS OF P450 CONTENT, GLUTATHIONE CONTENT (GSH), GLUTATHIONE-S-TRANSFERASE ACTIVITY (GST), AND P-AMINOPHENOL HYDROXYLASE ACTIVITY (PAP).

RESULTS

HIGH-DOSE DOGS OF BOTH SEXES AND MID-DOSE FEMALES DISPLAYED STATISTICALLY SIGNIFICANT INCREASES IN CYTOCHROME P450 (SEE ATTACHED SPECIAL LIVER CHEMISTRY TABLE). MEASUREMENT OF PAP INDICATED NO INDUCTION OF ACTIVITY WITH INCREASING DOSE (ALTHOUGH THE LOW-DOSE FEMALE VALUE WAS SIGNIFICANTLY ELEVATED ABOVE CONTROL VALUE BY THE DUNNETT'S T-TEST PERFORMED BY THIS REVIEWER). THE AUTHOR INDICATED THAT SINCE PAP SHOWS A SLIGHTLY GREATER SPECIFICITY OF INDUCTION POTENTIAL WITH P448 INDUCERS, THE DATA SUGGEST THAT THE TEST MATERIAL IS A P450 INDUCER AND NOT A P448 INDUCER (OFTEN A MARKER OF CARCINOGENIC POTENTIAL). THERE WAS NO DOSE-RELATED EFFECT REPORTED FOR EITHER SEX ON GSH OR GST, ALTHOUGH STATISTICAL SIGNIFICANCE WAS REPORTED IN THE MID-DOSE ANIMALS OF BOTH SEXES FOR GSH. THIS REVIEWER FOUND A DOSE-RELATED DECREASE (P<0.05) IN GST VALUES IN THE MID- AND HIGH-DOSE FEMALES USING DUNNETT'S T-TEST.

HISTOPATHOLOGY

THE FOLLOWING CHECKED (X) TISSUES/ORGANS WERE COLLECTED FROM ALL ANIMALS.

X HEART* BONE MARROW* X LYMPH NODES* X SPLEEN* X THYMUS* UROGENITAL X KIDNEYS*† X URINARY BLADDER* X TESTES*† X EPIDIDYMIDES X PROSTATE X SEMINAL VESICLE X OVARIES*† X UTERUS* CERVIX	X BRAIN*T X PERIPH. NERVE* (SCIATIC) X SPINAL CORD (3 LEVELS)* X PITUITARY* X EYES (OPTIC N.)* GLANDULAR X ADRENALS* LACRIMAL GLAND X MAMMARY GLAND* X PARATHYROIDS*TT X THYROIDS*TT OTHER X BONE* (STERNUM/FEMUR) X SKELETAL MUSCLE* X SKIN* X ALL GROSS LESIONS
	BONE MARROW* X LYMPH NODES* X SPLEEN* X THYMUS* UROGENITAL X KIDNEYS*† X URINARY BLADDER* X TESTES*† X EPIDIDYMIDES X PROSTATE X SEMINAL VESICLE X OVARIES*† X UTERUS* CERVIX

* REQUIRED FOR CHRONIC STUDIES

[†] ORGAN WEIGHTS REQUIRED IN CHRONIC STUDIES/TT FOR NON-RODENT STUDIES

RESULTS

THE LIVER WAS SHOWN TO BE THE TARGET ORGAN. LAMINAR EOSINOPHILIC INTRAHEPATO-CYTIC BODIES WERE OBSERVED IN ALL HIGH-DOSE MALES AND IN ONE MID-DOSE MALE. TWO HIGH-DOSE FEMALES ALSO DISPLAYED THESE BODIES. TWO HIGH-DOSE MALES DISPLAYED CANALICULAR BILE PLUGS, AND ONLY TREATED MALES SHOWED AN INCREASE IN INTRAHEPATIC PIGMENT.

OTHER OBSERVATIONS INCLUDE ONE HIGH-DOSE FEMALE WITH IMMATURE OVARIES AND ONE MID-DOSE MALE WITH DEGENERATION OF THE TESTICULAR GERMINAL EPITHELIUM. SINCE THIS FEMALE HAD NOT SHOWN ANY SIGN OF ESTROUS CYCLING DURING THE STUDY, THE AUTHOR STATED THAT IT WOULD BE IMPRUDENT TO DISMISS THE POSSIBILITY OF THIS FINDING BEING A CONSEQUENCE OF STEROID BIOSYNTHESIS INHIBITION.

CUNCLUS ION

THERE WERE NO DIFFERENCES OBSERVED IN SURVIVAL, OPHTHALMOSCOPIC PARAMETERS, OR HEMATOLOGIC PARAMETERS FOLLOWING THE ADMINISTRATION (DIET) OF SAN 619 F TO BEAGLE DOGS FOR 52 WEEKS AT DOSE LEVELS OF 30, 100, AND 350 PPM [1.0, 3.2, AND 12.1 (MALES); 12.6 (FEMALES) MG/KG/DAY, RESPECTIVELY].

BODY WEIGHT GAIN WAS LOWER IN THE HIGH-DOSE ANIMALS COMPARED WITH CONTROL, ALTHOUGH THE MAGNITUDE OF THE DIFFERENCE WAS NOT GREAT. THERE WERE DOSE-RELATED INCREASES IN PLATELETS THROUGHOUT THE STUDY IN BOTH SEXES, ALTHOUGH STATISTICAL SIGNIFICANCE WAS NOT ALWAYS ATTAINED. PROTHROMBIN TIMES WERE CHECKED AT WEEK 48 AND WERE SLIGHTLY ELEVATED IN THE MID-DOSE MALES AND THE HIGH-DOSE ANIMALS.

DIFFERENCES OBSERVED IN SEVERAL CLINICAL LABORATORY PARAMETERS BETWEEN THE CONTROL AND TREATED ANIMALS ARE CONSISTENT WITH EFFECTS ON THE LIVER (ELEVATED ALKALINE PHOSPHATASE AND ALAT LEVELS; DECREASED TOTAL PROTEIN, ALBUMIN, AND CHOLESTEROL LEVELS).

ABSOLUTE AND RELATIVE LIVER WEIGHTS WERE INCREASED IN THE HIGH-DOSE ANIMALS OF BOTH SEXES COMPARED TO CONTROLS, BUT STATISTICAL SIGNIFICANCE WAS ATTAINED ONLY IN THE MALES. RELATIVE KIDNEY WEIGHT WAS INCREASED (SIGNIFICANTLY) IN BOTH THE LOW- AND HIGH-DOSE FEMALES.

STATISTICALLY SIGNIFICANT INCREASES WERE OBSERVED IN CYTOCHROME P450 IN BOTH SEXES OF THE HIGH-DOSE AND IN THE MID-DOSE FEMALES. LAMINAR EOSINOPHILIC INTRAHEPATOCYTIC BODIES WERE OBSERVED IN ALL HIGH-DOSE MALES, ONE MID-DOSE MALE, AND TWO HIGH-DOSE FEMALES AND WERE CONSIDERED TO REPRESENT ADAPTIVE HYPERTROPHY OF THE ENDOPLASMIC RETICULUM.

THE NUEL CAN BE SET AT 30 PPM (1.0~Mg/kg/day) AND THE LEL AT 100~PPM (3.2~Mg/kg/day), BASED ON LIVER EFFECTS.

CYPROCONAZULE Tox review 007871
Page is not included in this copy. Pages 14 through 16 are not included.
The material not included contains the following type of information:
Identity of product inert ingredients.
Identity of product impurities.
Description of the product manufacturing process.
Description of quality control procedures.
Identity of the source of product ingredients.
Sales or other commercial/financial information.
X A draft product label.
The product confidential statement of formula.
Information about a pending registration action.
FIFRA registration data.
The document is a duplicate of page(s)
The document is not responsive to the request.
The information not included is generally considered confidential by product registrants. If you have any questions, please contact the individual who prepared the response to your request.

e en se